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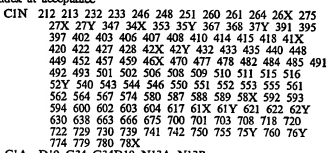
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(54) PREPARATION OF TITANIUM DIOXIDE CONCENTRATES AND OXIDE PIGMENTS FROM ILMENITE

We, BAYER AKTIENGESELLSCHAFT, formerly known as Farbenfabriken Bayer Aktiengesellschaft of 509 Leverkusen, Germany, a body corporate organised under the laws of Germany, do hereby declare the invention, for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following state-

This invention relates to a process for the production of titanium dioxide concentrates from ilmenite.

Processes for the production of titanium 15 dioxide concentrates containing 90% and more of TiO₂ from ilmenite are becoming increasingly important. These titanium dioxide concentrates are used instead of natural rutile, of which there are limited deposits, 20 for the production of titanium tetrachloride. Titanium tetrachloride is the starting material for the production of titanium dioxide pigments by the so-called chloride process.

The direct chlorination of ilmenite is uneconomical because during chlorination the iron present in the ilmenite (in quantities of from 30 to 40%) is converted into iron chloride. The amount of chlorine set aside for the iron is lost or has to be recovered

by complicated measures.

In one known process for working up

ilmenite, the iron oxide component is initially converted as completely as possible into metallic iron by a reduction process. Hydrogen, carbon monoxide, mixtures of both gases or solid carbon are mainly used as the reducing agents. Reduction can be carried out in rotary tubular kilns or other suitable apparatus such as fluidized beds or combustion chambers. The reaction temperatures are in the range of from 700°C. to 1200°C. The original shape of the ilmenite grain remains intact although, in addition to rutile, finely divided metallic iron is now present in the grain. Accordingly, it is not possible magnetically to separate the iron from the non-magnetic titanium dioxide, with the result that the iron component has to be dissolved out in a subsequent process with dilute acids or salt solutions. A high-percentage titanium dioxide concentrate is obtained which is suitable for use as a starting material for the production of titanium tetrachloride, cf., for example, British Patent Specification No. 1,104,262. In this known process, however, the iron accumulates in a form in which it

is almost impossible to use economically.

In the process disclosed in U.S. Patent Specification No. 3,252,787, the reduced ilmenite is treated with iron (III) chloride solution, giving a solid TiO₂-concentrate in addition to an iron(II)-chloride solution.

The iron(II)-chloride solution is then treated with atmospheric xygen to give iron(III)-chloride in addition to Fe_2O_3 . The iron oxide accumulates in the form of β -FeOOH and because of its high chloride content cannot be used as a pigment.

British Patent Specification No. 980,859 discloses a process in which, after a reducing roasting, the ilmenite is treated in an aqueous, preferably acid, medium, as a result of which the iron is converted into iron oxide while air or other oxidizing gases are blown in. Yellow or black oxides can be obtained, depending on the conditions.

15 However, considerable difficulties are involved in separating off the iron oxides from the TiO₂-concentrate. With the normal decantation process, the iron oxide contains up to 8% of TiO₂. It is also not possible with more sophisticated techniques of the kind known from the working up of ores considerably to reduce the titanium dioxide content of the iron oxide. According to Austral. Inst. Mining Metallurgy Proc. 214 25 (1965), page 42, it is unlikely ever to be possible to reduce the TiO₂ content to below 2%. However, iron oxides with a TiO₂ content such as this cannot be used as high-grade pigments.

It is also known that iron oxide pigments can be obtained by oxidizing iron and at the same reducing aromatic nitro com-pounds. The iron is reacted in a relatively coarse form (filings, wire tacks, etc.) in the presence of iron salts or even other metal salt solutions to form the required oxide. The starting material must have an iron content of at least 90%. Black Fe₂O₄ is obtained in the presence of iron salts, whilst yellow FeOOH is obtained in the presence of aluminium salt solutions. Red iron oxide pigments can be obtained by adding a separately prepared seed or nucleus suspension as known per se. The properties of the iron oxides formed can also be controlled as already known by other additives such as, for example, sulphuric acid, phosphoric acid or an aniline salt. Processes of this kind are described for example in German

Patent Specifications Nos. 463,773; 464,561 and 551,258. We have sought to provide a simple, effective and economic process for producing from ilmenite substantially iron-free 55 titanium dioxide suited for chlorination, and substantially titanium-free iron oxide pig-

In accordance with the present invention we provide a process for the production of 60 a titanium dioxide concentrate and an iron oxide pigment from ilmenite, in which ilmenite is subjected to heating under reducing conditions to reduce the iron content to metallic iron; the metallic iron in the reduced ilmenite is reacted with nitrobenz-

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cne at a temperature of from 20°C. to 100°C. and at a pH value of from 3 to 7 to form iron xide, and the iron oxide is separated from the titanium dioxide concentrate.

Surprisingly, the nitrobenzene reduction process gives an iron oxide pigment which can readily be separated off from the titanium dioxide residue. Not only is a highly concentrated titanium dioxide with a high TiO2 content of around 90% obtained, but an iron oxide whose TiO2 content is well below 1% is also obtained. Separation does not involve any complicated operations. It is possible to use the conventional grading apparatus such as, for example, screen graders and hydrocyclones, optionally in conjunction with overflow thickeners. Surprisingly, it is possible to obtain iron oxides with pigment properties which could not be obtained by the conventional nitrobenzene process. The black pigments are up to 80% superior in their tinting strength to the products made with iron filings by the conventional nitrobenzene reduction process. For the same tinting strength, the hue is tinged with much more blue than in conventional types. The yellow pigments are distinctly purer and are tinged with more green both in their full colour and also in admixture with titanium dioxide than the corresponding products made with iron filings with the aid of nitrobenzene. In some cases, it is even possible to obtain yellow pigments whose properties corres- 100 pond substantially to those of the pigment produced by the Penniman process. In the production of red pigments via a

seed suspension, long reaction times and an incomplete reaction are involved where the 105 usual iron filings or pins are used. By contrast, it is possible to obtain a complete reaction over a short period by using the reduced ilmenite.

The titanium dioxide concentrate separ- 110 ated off can be directly subjected to chlorination, although it is best treated with acids beforehand. In this way, it is possible to obtain concentrates containing 95% of TiO₂. After-treatment is carried out with the usual 115 inorganic acids such as, for example, hy-drochloric acid or sulfuric acid with concentrations of from 5 to 20% by weight in the case of hydrochloric acid and from 5 to 40% by weight in the case of sulphuric acid. It 120 is best to use the approximately 20% waste acid which is obtained from the TiO₂sulphate process and to carry out this process in several stages.

If the titanium dioxide concentrate and 125 the iron oxide pigment are to be cleanly separated from one another after the reaction with nitrobenzene, the reduced ilmenite used should not contain any fractions with grain sizes of less than 40 microns.

In a particular embodiment, the process according to the invention is carried out

The ilmenite to be reduced with particle sizes of from 40 to 200 microns is preferably initially subjected to an oxidation because a pre-oxidised ilmenite is casier to reduce. The oxidising pretreatment is carried out at a temperature of from 800 to 1100°C. in conventional apparatus such as for example a rotary tubular kiln or a fluidising furnace. The preoxidized ilmenite is then reduced, again in a rotary kiln, shaft furnace or fluidising furnace, under the 15 effect of a reducing agents at a temperature of from 700°C, to 1200°C. Preferred reducing agents include hydrogen, carbon monoxide or mixtures of both gases, or even natural gas, although it is also possible to 20 use solid carbon-containing materials such as, for example, coal dust. After cooling, the reduced ilmenite is transferred to vessels equipped with stirrers, in which the metallic iron is oxidized into iron oxide pigment following the addition of nitrobenzene and optionally the other conventional modifying compounds. On completion of oxidation, most of the crude aniline is decanted off from the paste-like reaction medium, and the residual aniline is separated off by distillation with steam. The aqueous suspension is then delivered to the separation stage for which it is possible to use any separation equipment by which fine-grained solids can be separated from one another through differences in specific gravity and grain size. As already mentioned, it is possible to use screen graders, overflow thickeners, hydrocyclones and other decanters, optionally in conjunction with one another. The titanium dioxide concentrate separated off is then preferably delivered to a leaching stage in which the titanium dioxide is treated with dilute inorganic acids, preferably in several stages. Most of the iron oxide still present is separated off during this after-treatment so that concentrates with a TiO₂-content of 95% are obtained in this way. Leaching is carried out in known manner in stirrer tanks or countercurrent apparatus. leaching, the solid residue is washed and dried and can then be subjected to chlorination. The iron oxide pigment separated off is also washed and dried and optionally

> The nitrobenzene reduction is carried out in known manner, the properties of the pigment being influenced by establishing special conditions.

In particular, the nitrobenzene reduction can be carried out as follows:

55 fine-ground.

FeCl₂, AlCl₃ and aniline salt solution are initially vigorously stirred either separately or together in different quantities and 65 ratios with some of the reduced ilmenite to

be treated and with some of the nitrobenzene to be used. The mixture very quickly reaches a pH of 3.5 to 4 and a temperature of 100°C, under the effect of the heat generated during the reaction. At this point, further quantities of reduced ilmenite and nitrobenzene are introduced and the mixture is kept at boiling temperature by the heat of recation until the required iron oxide has been formed at a pH of from 5 to 6.

The different yellow and black nuances of the iron oxide are obtained partly by the addition of relatively large quantities of FeCl₂, AlCl₃ and aniline salt solution and partly by the addition of relatively small quantities of H2SO4 and/or H2PO4

The process according to the invention is illustrated by the following Examples:

Example 1a-partially reduced ilmenite 1000 g. of ilmenite containing 50.0% of TiO₂, 37.7% of FeO and 9.1% of Fe₂O₃ with grain sizes of from 60 to 200 microns were initially oxidised with air for 1.5 hours at 800°C. in a fluidised bed. The subsequent reduction stage was also carried out at 800°C, with hydrogen over a period of 5 hours. The reduced ilmenite contained 40.5% of metallic iron.

Example 1b—black pigment
20 ml. of a 36.7% FeCl₂-solution (d.= 1.300), 20 ml. of water, 20 ml. of nitrobenzene and 200 g. of the partially reduced ilmenite containing 40.5% (81 g.) of Fe 100 metal and 56.2% of TiO2 are added to 90 ml. of a 43% aniline hydrochloride solution (d.=1.085). After this mixture has been heated while stirring to 90%°C. at a pH of 4, 105 ml. of nitrobenzene and 170 g. of 105 the partially reduced ilmenite containing 40.5 (69 g.) of Fe metal are simultaneously run in over periods of 2 hours and 1 hour, respectively.

On completion of the nitrobenzene re- 110 duction, most of the aniline formed is decanted off and the residue left in the reaction mixture is distilled off with steam.

The Fe₃O₄ formed by oxidation of the ilmenite iron is separated from the TiO₂- 115 concentrate by washing out, washed free from salts, filtered and dried at from 100°C. to 110°C. An iron oxide black pigment of high covering power coupled with a high resistance to light and alkalis is obtained 120 in this way. It has a particle size of from 0.5 to 0.6 micron and an Fe₃O₄-content (calculated at Fe₂O₂) of from 93 to 94%. Yield: more than 90%.

The ilmenite residue contained 91.1% of 125 TiO₂. After three leachings with hot 20% HCl (ratio by weight of concentrate to acid =1:1.2), a concentrate containing 94.1% of TiO₂ was obtained.

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Example 1c

In a rerun of Example 1b, the acid leaching was carried out with a waste acid, containing 20% of H₂SO₄, from the TiO₂ sulphate process (100 g. of residue to 100 ml. of acid) ver a period f 2 hours at 100°C. The moist residue was then heated once again with fresh acid for a period of 2 hours. The filtered and dried concentrate contained 95.2% of TiO₂.

Example 2—yellow pigment 60 ml. of water, 25 ml. of nitrobenzene and 63 g. of the partially reduced ilmenite 15 according to Example 1a, containing 40.5% (25 g.) of Fe metal are added to 80 ml. of 17.5% AlCl₂ solution (d.=1.205).

After this mixture has been heated with strring to 90°C. at a pH of 4.5, 142 ml. of nitrobenzene and 407 g. of the partially reduced ilmenite containing 40.5% (165 g.) of Fe metal are run in simultaneously over periods of 2 hours and 3 hours, respectively. In addition, approximately 300 to 400 ml. 25 of water are gradually run in after the reaction mixture-has assumed a certain consistency, in order to obtain the required consistency.

On completion of the nitrobenzene reduction, most of the aniline formed is decanted off and the residue left in the reaction mixture is distilled off with steam.

The FeO(OH) formed by oxidation of the ilmenite iron is washed out of the TiO: concentrate by washing, subjected to fur-ther washing to free it from salts, filtered and dried at a temperature below 200°C. An iron oxide yellow pigment of high covering power coupled with a high resistance to light and alkalis is obtained in this way. Its particle size is in the range of from 0.7 to 1.0 microns (length) and 0.2 micron wide, and its Fe₂O₃ content is in the range of from 86% to 88%. Oil number: 37, yield: more than 90%.

The residue freed from the finely divided pigment contained 86.1% of TiO2. Following the procedure described in Example 1b a concentrate containing 94.4% of TiO2 was 50 obtained with 20% HCl, whilst a concentrate containing 93.1% of TiO₂ was obtained with waste acid from the TiO2-sulfate process which contained approximately 20%

of free H2SO4.

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Example 3-red pigment

65 g. of reduced ilmenite, 416 ml. of FeSO, (20%) and 100 ml. of nitrobenzene are adjusted to pH 8.6 with 210 ml. of 19% 60 NaOH. 162 ml. of nitrobenzene and 790 g. of partially reduced ilmenite are run into

this seed solution at 100°C. ver periods of 1 hour and 4 hours, respectively, such that the pH is 4-5. The reaction is complete after 5 hours. A red pigment is obtained from the reaction mixture which may be further worked up as in Examples 1

WHAT WE CLAIM IS:-

1. A process for the production of a titanium dioxide concentrate and an iron oxide pigment from ilmenite, in which ilmenite is subjected to heating under reducing conditions to reduce the iron content 75 to metallic iron; the metallic iron in the reduced ilmenite is reacted with nitrobenzene at a temperature of from 20°C, to 100°C. and at a pH value of from 3 to 7 to form iron oxide; and the iron oxide is separated from the titanium dioxide concentrate.

A process as claimed in claim 1, in which the reaction with nitrobenzene is carried out in the presence of a hydrolysable metal salt, a seed or other modifying compound.

3. A process as claimed in claim 1 or 2. wherein a black pigment is prepared in the presence of an iron salt.

4. A process as claimed in claim 1 or 2, wherein a yellow pigment is prepared in the presence of an aluminium salt.

5. A process as claimed in claim 1 or 2, wherein a red pigment is prepared by adding an Fc(OH)₂ seed suspension.

6. A process as claimed in any of claims 1 to 5, wherein the separated titanium dioxide concentrate is treated with an acid 100 remove residual iron.

7. A process as claimed in claim 6. wherein the approximately 20% waste acid obtained from the TiO₂-sulphate process is used for the acid treatment.

8. A process as claimed in any of claims 105 1 to 7, wherein the reduced ilmenite does not contain any fractions with grain sizes of less than 40 microns.

9. A process according to claim 1, substantially as herein described with reference 110 to any of the specific Examples.

10. A titanium dioxide concentrate and an iron oxide pigment, when made by a process according to any of claims 1 to 9.

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